

MICROCOPY RESOLUTION TEST CHART NATIONAL BUREAU OF STANDARDS-1963-A



OFFICE OF NAVAL RESEARCH Contract NO0014-79-C-0670

TECHNICAL REPORT NO. 34

Surface Environmental Effects in Electrochemical
Kinetics: Outer-Sphere Chromium(III) Reductions at
Mercury, Gallium, Lead, and Thallium Surfaces

Ъу

H. Y. Liu, J. T. Hupp, M. J. Weaver

Prepared for Publication

in the

Journal of Electroanalytical Chemistry

Department of Chemistry

Purdue University

West Lafayette, IN 47907

September 1984



Reproduction in whole or in part is permitted for any purpose of the United States Government

This document has been approved for public release and sale; its distribution is unlimited

84 09 18 247

REPORT DOCUMENTATION	READ INSTRUCTIONS BEFORE COMPLETING FORM		
REPORT NUMBER	2. GOVT ACCESSION NO.	3. RECIPIENT'S CATALOG NUMBER	
Technical Report No. 34			
TITLE (and Subtitle)		5. TYPE OF REPORT & PERIOD COVERER	
Surface Environmental Effects in Electrochemical Finetics: Outer-Sphere Chromium(III) Reductions at Mercury, Gallium, Lead, and Thallium Surfaces		Technical Report No. 33 6. PERFORMING ORG. REPORT NUMBER	
H. Y. Liu, J. T. Hupp, M. J. Was	aver	x00014-79-C-0670	
PERFORMING ORGANIZATION NAME AND ADDR Department of Chemistry Purdue University West Lafayette, IN 47907	ESS	10. PROGRAM ELEMENT, PROJECT, TASK AREA & WORK UNIT NUMBERS	
Controlling office name and address Office of Naval Research		12. REPORT DATE September 1984	
Department of the Navy Arlington, VA 22217		13. NUMBER OF PAGES	
4. MONITORING AGENCY NAME & ADDRESS(II dill	ferent from Controlling Office)	15. SECURITY CLASS. (of this report)	
		Unclassified	
		154. DECLASSIFICATION/DOWNGRADING	
Approved for Public Release; d		<u> </u>	

DISTRIBUTION ST. 4ENT (of the abstract entered in Block 20, if different from Report)

.E. SUPPLEMENTARY .CTES

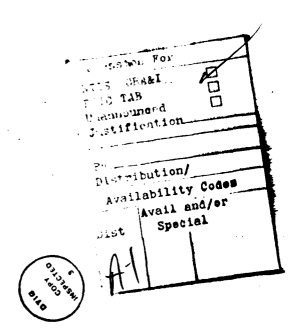
'S. KEY WORDS (Continue on reverse eide if necessary and identify by block number)

underpotential deposited (upd); rate alterations; durface hydrophilicity; nonadiabatic reaction pathways

ABSTRACT (Continue on reverse side if necessary and identify by block number)

Electrochemical rate parameters for the outer-sphere reduction of several Cr(III) aquo and ammine complexes and also for $\text{V}(\text{OH}_2)_6^{-3+}$ and $\text{Eu}(\text{OH}_2)_1^{-3+}$ are compared at aqueous-metal interfaces formed with mercury, liquid gallium, lead, and underpotential deposited (upd) lead and thallium monolayers on silver. For reactants containing aquo ligands, substantial (up to 10^3 -fold) decreases in the rate constants, both before and after electrostatic double-layer corrections, were observed at a given electrode potential when substituting

mercury by the other surfaces, especially lead and gallium. alterations are accompanied by marked decreases in the apparent activation entropies, although offset by corresponding decreases in the measured activation enthalpies. These results are interepreted in terms of the varying influence of these metal surfaces on the interfacial solvent structure. The observed substrate dependence Hg > upd Pb ~ upd Tl ≥ Pb > Ga is consistent with the anticipated differences in surface hydrophilicity. This suggests that the effects arise either from modifications to the solvation energetics of the outer-sphere transition state, or from differences in the double-layer potenital profile caused by water dipole orientation. The likely influence of nonadiabatic reaction pathways, is also considered. Smaller rate variations were observed for Cr(NH₃)6 and reduction (en = ethylenediamine), although the activation parameters are more sensitive to the metal substrate. The relatively small influence exerted by mercury surfaces upon the outer-sphere reaction energetics is also consistent with the reasonable agreement seen between the experimental and theoretical rate parameters for Cr(OH2)6 at this surface.



SURFACE ENVIRONMENTAL EFFECTS IN ELECTROCHEMICAL KINETICS: OUTER-SPHERE CHROMIUM(III) REDUCTIONS AT MERCURY, GALLIUM, LEAD, AND THALLIUM SURFACES

H. Y. Liu[†], Joseph T. Hupp and Michael J. Weaver Department of Chemistry, Purdue University
West Lafayette, IN 47907 U.S.A.

vanadium and suresium

ABSTRACT

> Electrochemical rate parameters for the outer-sphere reduction of several Cr(III) ague and ammine complexes and also for $(V(OH_2))_6$ and Eu(OH₂) compared at aqueous-metal interfaces formed with mercury, liquid gallium, lead, and underpotential deposited (upd) lead and thallium monolayers on silver. For reactants containing aquo ligands, substantial (up to 103-fold) decreases in the rate constants, both before and after electrostatic double-layer corrections, were observed at a given electrode potential when substituting mercury by the other surfaces, especially lead and gallium. The rate alterations are accompanied by marked decreases in the apparent activation entropies, although offset by corresponding decreases in the measured activation enthalpies. These results are interpreted in terms of the varying influence of these metal surfaces on the interfacial solvent structure. The observed substrate dependence Hg > upd Pb ~ upd T1 > Pb > Ga is consistent with the anticipated differences in surface hydrophilicity. This suggests that the effects arise either from modifications to the solvation energetics of the outer-sphere transition state, or from differences in the double-layer potential profile caused by water dipole orientation. The likely influence of nonadiabatic reaction pathways is also considered. Smaller rate variations were observed for $Cr(NH_3)_6^{3+}$ and $Cr(en)_3^{3+}$ reduction (en = ethylenediamine), although the activation parameters are more sensitive to the metal substrate. relatively small influence exerted by mercury surfaces upon the outer-sphere reaction energetics is also consistent with the reasonable agreement seen between the experimental and theoretical rate parameters for $Cr(OH_2)_6^{3+}$ reduction at this surface.

Graduate Research Assistant, Michigan State University, 1979-82.

Author to whom correspondence should be addressed.

INTRODUCTION AND CONCEPTUAL BACKGROUND

A central fundamental question in electron-transfer kinetics at metal-solution interfaces concerns the manner and the extent to which the chemical nature of the metal surface influences the overall reaction energetics. The issues involved can usefully be perceived in terms of the following rate expression 1,2:

$$k_{app} = K_p \kappa_{el} \Gamma_n \nu_n \exp(-\Delta G^*/RT)$$
 (1)

where K_p is the equilibrium constant (cm) associated with transporting the reactant from the bulk solution to the reaction site (the "precursor state"), ν_n is the nuclear frequency factor (sec⁻¹), κ_{el} is the electronic transmission coefficient, Γ_n is a nuclear tunneling factor, and ΔG^* is the free energy of activation. The "encounter preequilibrium" rate formalism embodied in Eq. (1) has been described in detail¹; it is physically more realistic than the conventional "collision" treatment. Thus the former model more correctly treats the overall reaction as a two-step process involving the unimolecular activation of reactant within a precursor state that is in quasi-equilibrium with respect to the bulk reactant state. \frac{1}{2}

Equation (1) is most obviously applicable to inner-sphere reaction pathways, i.e. where the reactant is bound directly to the electrode surface in the precursor state. However it is also applicable to outer-sphere pathways, i.e. where the reactant undergoes electron transfer without penetrating the inner solvent layer. The chemical nature of the metal surface is unquestionably important for inner-sphere processes since K_p and possibly ΔG^* and κ_{el} can be influenced strongly by the specific reactant-surface forces involved. However, the influence of the metal surface upon the energetics of outer-sphere reactions is rather more subtle, and remains a controversial topic. We present here new experimental data aimed at provicing unabmigious information on this question for some reactions at metal-aqueous interfaces. In order to clarify the approach taken, a brief description of the underlying physical models will first be presented.

-

≥ sc-called "weak-overlap" limit may usefully be envisaged for outer-sphere TETTES where the precursor state reactant interacts sufficiently weakly with the ======= so that the reorganization energy AG* will be essentially unaffected by the This will clearly be the case for reaction sites Interest some distance from the surface. However such sites are not expected to minimum importantly to the measured rate since the electronic coupling between the THETHE SUTFace and reactant orbitals will likely be insufficient to yield significantl \rightarrow values of κ_{el} . This electronic coupling will progressively increase as the Teactant-electrode distance, r, decreases, eventually yielding "adiabatic pathways"; === == κ_0 = 1. This "electronic factor" will thereby strongly favor reaction mites close to the surface, to an extent determined by the dependence of κ_{al} on r. Exercipe such sites may well be associated not only with significantly different -=-= $cf \Delta G^*$, but also with different values of $K_{_{\mathbf{D}}}$ as a result of the alterations The surface environment of the reactant caused by its proximity to the metal surface The swerall measured rates will therefore arise from an integral of "local" TELES associated with various reaction sites, appropriately weighted according to the $\kappa_{\rm el}$, $\kappa_{\rm p}$ and $\kappa_{\rm el}$, $\kappa_{\rm p}$ and $\kappa_{\rm el}$ associated with each site. mer therefore contain dominant contributions from sites sufficiently close to the Electrode so to yield rate parameters that are sensitive to the local surface environ substantially different from those expected on the basis of the weak overlap mode

Terms of Coulombic double-layer effects. Thus according to the Frumkin relation are agreent rate constant at a given electrode potential, k_{app}^{E} , can be expressed as

$$\log k_{app}^{E} = \log k_{corr}^{E} - (F/2.303 RT)(Z_{r} - \alpha_{corr})\phi_{r}$$
 (2)

in the reactant charge number, ϕ_r is the average electrostatic potential in the reaction site, and k_{corr}^E and α_{corr} are the work-corrected rate constant and

transfer coefficient, respectively, at the same electrode potential, E. This effect can be incorporated into the above preequilibrium rate formalism by rewriting Eq. (1) as

$$k_{corr} = K_{corr} \Gamma_{n} \nu_{exp} \left(-\Delta G_{corr}^{*}/RT\right)$$
 (3)

where
$$K_o = K_p \exp(-Z_r F \phi_r / RT)$$
 (4)

and
$$\Delta G^* = \Delta G^* - \alpha_{corr}^{F\phi}$$
 (5)

Equations (4) and (5) contain the components of the electrostatic double-layer correction in Eq. (2) associated with the formation of the precursor state and with the elementary electron-transfer step, respectively.

It is often presumed that the influence of the metal surface upon k_{app} is wholly described by Eq. (2). This notion has been fostered by the relative success of such relations in describing double-layer effects at mercury electrodes. Enwayer, there are good reasons to doubt this. Besides the niceties of discreteness-of-charge effects, the solvation environment at the reaction site may differ significantly from that in the bulk solution, thereby influencing K_p and ΔG_{corr}^* in a manner beyond that described by Eqs. (4) and (5). In addition, it is possible that K_{corr}^* << 1 even at the plane of closest approach, yielding overall "nonadiabatic" reaction

^{*}Two such effects can be distinguished in the absence of specific ionic adsorption of the supporting electrolyte. The "self-image" energy is the effect upon ϕ_r , and hence the work terms, arising from the attraction between the reacting ion and its electrostatic image in the metal. This effect is probably negligible for outer-sphere reactions. However, another component arises from the effect upon ΔG^*_{corr} from the attraction between the transition-state species and its image in the metal. The latter, which is included in Marcus' theoretical treatment, cannot be wholly neglected since the nonequilibrium polarization in the transition state prevents diffuse-layer screening from being entirely effective. Nevertheless, its magnitude is estimated to be small or moderate for outer-sphere reactions (ca 1-1.5 k cal mol -1) and essentially independent of the electrode material.

pathways. Such factors are unlikely to be exposed by rate measurements at a single metal surface since they are liable to remain approximately constant under these conditions.

A stringent test of these possibilities would be to examine the effects of altering the chemical nature of the electrode material upon k_{corr}^{E} for well-defined outer-sphere reactions, preferably for surfaces known to exert disparate influences upon the local solvent structure. Few such examinations have been made. We have referred to most of these studies in an earlier article on this topic. A brief review is also available. Although several systems, such as benzoquinone reduction in dimethylformamide and some anion reductions in aqueous media yield values of k_{corr} that are approximately independent of the electrode material, others show large differences. We have found that the electrooxidation kinetics of some aquo complexes are substantially slower at platinum and gold relative to mercury electrodes, although the electroreduction of Co(III) ammines yields similar rate parameters at these surfaces.

The "double-layer corrections" [Eq. (2)], can be varied systematically by altering the coordinated ligands. The standard potentials, and hence the standard rate the standard rate that is, for some of these reactions are unknown on account of their chemical irreversibility. All that is required, however, is the comparison of rate that it is that is required, however, is the comparison of rate that it is at different surfaces at a fixed electrode potential, whereupon the (albeit unknown) overpotential for each reaction will remain constant.

We have selected metal surfaces for which compositional information could be obtained from differential capacitance-potential data so that values of ϕ_r can be calculated, and which exhibit sufficiently high hydrogen overpotentials to enable Ir(III) reduction kinetics to be studied over a reasonable potential range in weakly acidic media. Besides mercury, gallium and lead were found to be suitable. 23 in though silver exhibits insufficiently high hydrogen overpotentials, underpotential deposited (upd) monolayers of lead and thallium on silver were also found to be satisfactory. An additional reason for selecting the upd lead surface was to tempare its adsorptive and electrochemical kinetic properties with those of bulk lead electrodes. We have noted that these two surfaces have surprisingly similar facuble-layer properties. The metals are anticipated to bring about significantly different interfacial environments in aqueous media resulting from the decidely more "raydrophilic" nature of lead, thallium and especially gallium, in comparison with mercury. The comparison of gallium and mercury is of particular interest since the former can also be examined as a liquid close to room temperature (30°C).

Electrochemical rate constants and activation parameters are reported here for the reduction of six Cr(III) aquo and ammine complexes, along with $\mathrm{Eu}(\mathrm{OH}_2)^{3+}_n$ and $\mathrm{V}(\mathrm{OH}_2)^{3+}_6$ in aqueous solution at mercury, gallium, lead, and upd lead-silver and reallium-silver surfaces. Taken together, they illustrate the important influences that the interfacial solvation environment can exert on the reaction energetics for outer-sphere electrochemical reactions.

ENGERIMENTAL

Materials and Electrodes

The various Cr(III) aquo and ammine complexes employed here were synthesised using procedures outlined in refs. 16 and 18; $Cr(OH_2)_6^{3+}$, $Cr(OH_2)_5^{7+}$, and $Cr(OH_2)_5^{7+}$ could be isolated as the solid perchlorate salts. Stock solutions of $Eu(OH_2)_6^{7+}$ and $V(OH_2)_6^{7+}$ were prepared as in ref. 11. This paper also contains details of the supporting electrolyte preparation, etc.

A dropping mercury electrode (d.m.e.), having a flow rate around 2 mg sec-1 and a mechanically controlled drop time was used to examine electrode kinetics at this metal. Measurements at gallium (99.99%, Research Inorganic Co.) were performed in a thermostatically controlled glove box held at 30°C, using a commercial micrometercommodiled hanging drop electrode (Brinkman Instruments, Inc.). The gallium drops were used within a few secondsafter formation so to minimise the accumulation of trace impurities at the surface. The solid metal substrates used (lead and silver) were fabricated from high purity (99.999%) polycrystalline rods (Atomergic Chemical Co., Materials Research Corp.) as rotating disk electrodes, having a disk radius of 0.20 cm and a Teflon (or Kel-F) sheath radius of 0.6 cm. They were either purchased from Pine Instruments, or constructed in the department. The lead electrodes were fabricated by gluing the machined lead rod to the stainless steel support with conductive silver-filled expoxy (Transene Co.). They were then pressed into a hot Teflon or Kel-F sheath machined such that a leak-free fit was obtained upon cooling. 23 The silver electrodes were similarly prepared, the silver roc being soldered to the stainless steel shaft. 29 For temperature-dependent studies, electrodes were prepared that featured instead a spiral copper spring contacting the metal roc and the steel mount in order to minimise heat loss from the electrode surface. 29

Several methods were examined for pretreating the lead surfaces prior to the electrochemical measurements. In method A, 30 the surface was initially mechanically polished with 1 km alumina on a polishing wheel using water as a lubricant. The electrode was then rinsed a number of times with a 2:3:5 mixture of acetic acid, 30% hydrogen peroxide, and methanol. After a final rinse with water, the electrode was rapidly transferred while wet to a degassed 0.5 M sodium perchlorate solution and held at -1.5 V vs a saturated calomel electrode (s.c.e.) for thirty minutes. In method B, 31 the electrode was mechanically polished on roughened glass, and electrochemically etched in 20% perchloric acid. Both methods A and B, when applied with care, could yield reasonably reproducible surfaces exhibiting well-defined capacitance and electrochemical kinetic behavior 23 (vide infra). However, the most satisfactory results were generally obtained by polishing with 1 km alumina until a shiny surface was obtained, rapidly rinsing with degassed water, and transferred immediately to 0.5 M NaClo4, the electrode potential thereupon being scanned repeatedly between -C.7 V and -1.6 V vs s.c.e. (Method C).

The upd lead-silver and thallium-silver surfaces were prepared essentially as described for the former in ref. 25. This involved the underpotential deposition of lead or thallium on an electrochemically pretreated silver substrate using dilute (ca C.5 µM Pb²⁺ or Tl⁺) solutions, the rate of deposition being controlled by rotating the electrode. In the thallium case, this entailed using a potential around -C.96 V vs s.c.e., rotating at 600 r.p.m. until a thallium monolayer had been formed. This point was determined from the anodic charge required to remove the layer using linear sweep stripping voltammetry. Monolayer formation corresponded to the time required for successive deposition beyond which no further increases occurred in the anodic stripping peak. The upd surfaces were prepared in Situ, prior to satisfy the reactant species for which rate data were required. The use of such dilute Pb²⁺ or Tl⁺ solutions enabled the kinetic as well as capacitance properties

the electrode to be examined over a range of potentials negative of the deposition potential either by using quiescent solutions or short potential pulses (vide infra).

Techniques

All the present Cr(III) reduction reactions occur at sufficiently high cathodic exerpotentials to render negligible the influence of the reoxidation of Cr(II) upon the measured kinetics, even using "slow perturbation" techniques such as d.c. Follarography. 16,18 (This is a fortunate circumstance since, except for $Cr(OH_2)^{3+/2+}$ =11 the present Cr(III)/(II) reactions are chemically irreversible. 16,18) Most rate data at solid as well as mercury electrodes in the present study were obtained Er means of normal pulse polarography, using 1-2 mM bulk reactant concentrations. Values of k and as a function of potential were obtained from the Oldham-Parry amalysis. 32 At mercury, the pulses were synchronized with the d.m.e. using a PAR 174A Folarographic Analyzer (EG&G Inc.). At solid surfaces, the potential pulse train was applied while the electrode was rotated at about 700 r.p.m. This condition emabled the reactant to be entirely replenished within the diffusion layer during the 2 or 5 sec delay between pulses without significantly influencing the diffusion profile during the short (50 msec.) potential steps. Some measurements at lead also utilized conventional rotating-disk voltammetry. The analysis is described in ref. 11. Pulse polarography proved to be an efficacious technique for obtaining E - E data at the upd surfaces, since no further metal deposition was found to cocur during the polarographic pulses. This was true even when the potential was stepped to values where bulk lead or thallium deposition occurs, providing that the initial potential selected lay within the upd region. 24 Rate data at hanging gallium electrodes were obtained by linear sweep voltammetry and analyzed by means of the relations in ref. 33. Sweep rates from 50-500 mV sec⁻¹ were employed. This method as also employed at the other surfaces, good agreement generally raing obtained with the rate constants obtained from normal pulse polarography.

Electrochemical rate measurements, except at gallium, were made at 3-6 temperatures over the range 2°-40°C using the nonisothermal cell rangement described in ref. 19. This entailed holding the reference electrode at room temperature while the cell temperature was varied. Electrode pretreatment was required at each new temperature. Rate data at gallium were obtained at temperatures 2-5°C above its melting point (29°C) and extrapolated to 25°C in order to facilitate comparisons with data gathered at the other surfaces at this temperature.

Differential capacitance measurements employed a Wein bridge and null detection as described in ref. 34. Other experimental details are given in ref. 11. All potentials were measured and are quoted versus the s.c.e.

RESULTS AND DATA TREATMENT

Electrochemical Rate Constants: Double-Layer Corrections

Table I contains a summary of rate constants, k_{app} , obtained at -1000 mV for six Cr(III) complexes, and also Eu(OH₂) $_{n}^{3+}$ and V(OH₂) $_{6}^{3+}$, in aqueous solution at mercury, gallium, lead, upd lead and upd thallium surfaces at 25°C. The listed values of k_{app} were measured in 0.5 M NaClO₄ + 3mM HClO₄, except those for Cr(NH₃) $_{6}^{3+}$ reduction which were determined instead in 40 mM La(ClO₄) $_{3}$ + 3 mM HClO₄ due to the limited solubility of this complex in concentrated perchlorate media. Similar values of k_{app} were obtained for the other reactants if 40 mM La(ClO₄) $_{3}$ was substituted for 0.5 M NaClO₄. Where necessary, the values of k_{app} at -1000 mV were obtained by linearly extrapolating plots of log k_{app} against E; the measured values of k_{app} spanned the range ca 10⁻⁴ to 4 x 10⁻² cm sec⁻¹. The selection of -1000 mV as the common potential minimised the extent of such extrapolation. It is also sufficiently negative to largely avoid perchlorate specific adsorption (vide infra). Apparent transfer coefficients, α_{app} , are also listed in Table I. These were obtained from $\alpha_{app} = -(RT/F)(dlnk_{app}/dE)$. These values were essentially independent of potential within the ca 200-300 mV scential range for which sufficiently precise values of k_{app} could be obtained.

The values of k_{app} obtained at mercury were generally reproducible within at least L5-20%. Somewhat inferior reproducibility was encountered at liquid gallium, although this was usually within ca 50%, similar to that obtained at both upd lead and thallium surfaces. Somewhat greater difficulties were experienced at solid lead surfaces. The original pretreatment method of choice, Method A above, yielded irreproducible ralues of k_{corr} that rapidly decreased with time. Method B did not fare much better. Nevertheless, when applied diligently Method C yielded values of k_{app} that were stable within ca. twofold for considerable periods (20-30 mins) following surface preparation. This pretreatment also yielded surfaces exhibiting well-defined differential capacitance behavior. Since k_{app} obtained using Method A approached those found with Method C after about twenty minutes. Nevertheless, all these pretreatments exhibited virtually identical (± 0.01-0.02) values of α_{app} .

As a check on the possibility that the small values of k_{app} obtained at lead might be due to adsorption of trace organic contaminants, some kinetic experiments were performed with solutions containing 0.1 mm Pb²⁺. Continuous lead deposition occurs under these conditions, presumably renewing the surface at a faster rate than that of contaminant adsorption even if the latter is diffusion controlled. (We thank for this suggestion.) Essentially identical that parameters were obtained in the absence and presence of Pb²⁺.

Work-corrected rate constants, k_{corr} , at -1000 mV were determined from the Listed values of k_{app} by applying Eq. (2). The values of α_{corr} were obtained from the corresponding values of α_{app} as noted below. The required values of ϕ_r were fetermined for the complexes containing predominantly aquo ligands by assuming that $\epsilon_{\pm} \approx 0.6 \ \zeta_2$, where ϕ_2 is the Gouy-Chapman diffuse-layer potential determined from the corresponding electronic charge density, q^m . This assumption is supported by extensive kinetic data gathered for aquo reactants at mercury, and is consistent with their large hydrated radii. ¹⁴,21 For the less hydrated ammine reactants, it

assumed that $\phi_r \approx \phi_2$, again on the basis of extensive rate-double layer comparison are cury electrodes. The required values of q^m for mercury were taken from the layer comparison is need tables, and for gallium by integrating capacitance data from the known are calculated as a contract of zero charge (p.z.c.) in perchlorate media (-935 mV vs sce).

The q^E values at the three solid surfaces were obtained as follows. ^{23,24}

Elles of the p.z.c. were determined in sodium fluoride from the position of the

====citance-potential minimum in dilute electrolytes (0.01-0.05 M). (The use of

====citance-potential minimum in dilute electrolytes (0.01-0.05 M). (The use of

====citance media was necessitated by the occurrence of significant perchlorate specific

=====ption at the p.z.c., as evidenced by the broader and concentration-dependent

====citance minima obtained in dilute perchlorate electrolytes. ^{23,24}) Lead, upd lead

=====citance minima obtained in dilute perchlorate electrolytes. ^{23,24}) Lead, upd lead

=====citance minima obtained in dilute perchlorate media of -800, -800, and -960 mV,

=====citively. Combining these values with integrated double layer capacitance
=====citance minima obtained in dilute perchlorate double layer capacitance
=====citance minima obtained in dilute perchlorate media were obtained by back integrating

=====sponding q^m=E curves in perchlorate media were obtained by back integrating

=====sponding fluoride and perchlorate electrolytes will be coincident. ²⁴

This procedure yielded q^m values at -1000 mV at lead, upd lead-silver, and upd mathematical within experimental error, \pm 0.3 μ C cm⁻², for 0.5 M NaClO₄ and la(ClO₄)₃, respectively.) Slight perchlorate specific adsorption was detected at each surface at -1000 mV from an analysis of C_{dl} -E curves in mixed fluoride-surface electrolytes. Inclusion of the adsorbed perchlorate charge densities, in the defective inner-layer charge densities, (q^m + q'), and resulting ϕ_2 recently as follows: lead, (q^m + q') = -3.0 μ C cm⁻², ϕ_2 = -18 mV (0.5 M NaClO₄), lead-silver, (q^m + q') = -2.0 μ C cm⁻², ϕ_2 = -13 mV ϕ_2 NaClO₂, -13.5 mV (40 mM La(ClO₄)₃); upd thallium-silver, (q^m + q') = -2.0 ϕ_2 cm⁻², ϕ_3 = -13 mV ϕ_3 NaClO₄. [Values of (q^m + q') in 0.5 M NaClO₄

and 40 mM La(ClO $_4$) $_3$ were found to be identical within experimental error, \pm 0.5 mC cm $^{-2}$.] Full details of the double-layer capacitance measurements are given elsewhere. Representative ϕ_2 -E curves determined for upd lead-silver are shown in Fig. 1. Electrostatic double-layer corrections were also applied to the capp values to yield the corresponding work-corrected values, $\alpha_{\rm corr}$, (Table I) by using the relation 14

$$\alpha_{corr} = \{ [\alpha_{app} - Z_r(d\phi_r/dE)]/[1 - (d\phi_r/dE)] \}$$
 (6)

The required values of $(d\phi_r/dE)$ for the various surfaces within the potential region where α_{app} was evaluated were obtained from the integrated C_{dl} -E data using the above procedure; these are listed in the footnotes to Table I. Other pertinent details are given in refs. 14 and 17.

A key question concerns the possibility that the sizable differences in \boldsymbol{k} seen for most reactants at the different surfaces could be due primarily to systematic errors in the electrostatic double-layer corrections. However, two lines of evidence indicate that this is not the case. The first is obtained from rate data gathered for these reactants at mercury electrodes in a variety of electrolytes. The responses of k_{app} to systematic alterations in the double-layer structure are quantitatively consistent with Eqs. (2) and (6) once the different sizes of the supporting electrolyte cations forming the outer Helmholtz plane (o.H.p.), along with the likely deficiencies of the Gouy-Chapman model, are taken into account. $^{14-18,21}$ Bearing in mind the magnitude of these double-layer corrections for the electrolytes considered here the resulting uncertainties in $k_{\mbox{corr}}$ on this basis are unlikely to be greater than 2-5 fold for tripositive reactants, and less for those having smaller charges. The extent of such double-layer corrections at the other surfaces is small due to the proximity of the p.z.c. values to -1000 mV. This is supported by the chserved approximate independence (within ca 2-fold) of $k_{\scriptsize{\mbox{app}}}$ at these surfaces to the supporting electrolyte composition. Secondly, the dependence of $k_{\mbox{\footnotesize corr}}$ upon the

electrode material for $Cr(OH_2)_5F^{2+}$ and $Cr(OH_2)_5OSO_3^+$ reduction is comparable to, or even larger than, that seen for $Cr(OH_2)_6^{3+}$ reduction (Table I). This is opposite to the result expected if electrostatic double-layer effects were primarily responsible for these rate differences. Thus on the basis of Eq. (2), since $(Z_r - \alpha_{corr}) \approx 0.5$ for $Cr(OH_2)_5OSO_3^+$, and $(Z_r - \alpha_{corr}) \approx 2.5$ for $Cr(OH_2)_6^{3+}$, the latter reactant should show about 5-fold greater variations of $\log k$ than for the former, in complete contrast to the experimental results (Table I). This tactic of employing reactants containing nonadsorbing anionic ligands in order to alter Z systematically has been exploited previously to distinguish between metal substrate effects upon $k_{\scriptsize{\mbox{app}}}$ arising from electrostatic and from specific surface effects. 11,37 Its validity for the present systems is supported by the observation that the reductions of $Cr(OH_2)_5F^{2+}$ and $Cr(OH_2)_5OSO_3^+$ occur by outer-sphere pathways at mercury electrodes, their sensitivity to alterations in the double-layer structure being in close accordance with Eq. (2). 16,17 Outer-sphere pathways are also virtually certain at the other surfaces studied here in view of the virtual absence of F or 50_{λ}^{2-} specific adsorption in the potential region where the rate data were gathered. 23,24,38

Electrochemical Activation Parameters

Given the observed sensitivity of $k_{\rm corr}$ to the nature of the metal surface, it is of interest to explore how this dependence is reflected in the temperature dependence of $k_{\rm corr}$. We have demonstrated how the evaluation of electrochemical activation parameters can shed light on the nature of reactant-solvent interactions in the transition state for electron transfer. 1,10,19,20,39 Using the preequilibrium rate formalism [Eq. (1)] we can write 1

$$k_{corr} = \delta r \kappa_{el} \Gamma_n v_n \exp(\Delta S_i^*/R) \exp(-\Delta H_i^*/RT)$$
 (7)

where K_0 has been replaced with an effective "reaction zone thickness", δr , and ΔS_1^* and ΔH_1^* are the so-called "ideal" entropies and enthalpies of activation. 19,40 These activation parameters represent the actual entropic and enthalpic barriers to electron transfer at the particular potential at which they are evaluated; ΔH_1^* can be evaluated from the slope of an Arrhenius plot of Rlnk corr versus (1/T) at a constant nonisothermal cell potential. 19,40 Such temperature-dependent measurements of k corr therefore enable estimates of the combined preexponential factor $\delta r_{el} r_n r_n$ to be obtained if ΔS_1^* can be estimated or, conversely, enable ΔS_1^* to be determined if the value of preexponential factor is assumed.

Calculated values of ΔS_{i}^{*} , ΔS_{calc}^{*} , can be obtained from 19,40,41

$$\Delta S_{calc}^* = \Delta S_{int}^* + \alpha_{corr} \Delta S_{rc}^*$$
 (8)

where ΔS_{rc}° is the thermodynamic entropy difference between the reduced and oxidized forms of the redox couple (the "reaction entropy" $^{42-44}$), and ΔS_{int}^{\star} is the intrinsic activation entropy. It is important to note that ΔS_{int}^{\star} will be close to zero (within ca 15 J K⁻¹ mol⁻¹ for couples in aqueous media) even when specific solute-solvent interactions are taken into account, provided that the transition-state entropy is related quadratically (or more linearly) to the entropies of the reactants and product species in the bulk solution. However, this will be the case only if the solvation environment in the transition state is similar to that

in the bulk solution. Therefore experimental estimates of ΔS_1^* that differ substantially from $\Delta S_{\rm calc}^*$ provide evidence that the transition-state solvation is perturbed by the electrode surface. The identification of the Arrhenius slope with ΔH_1^* presumes that the various preexponential terms in Eq. (7) are temperature independent. It is therefore necessary to apply a correction to allow for the anticipated temperature dependence of Γ_n . Analytical expressions are available. Since Γ_n decreases with temperature, this will tend to decrease the overall preexponential factor. It is usual to treat this term as a component, ΔS_1^* , of $\Delta S_{\rm int}^*$, thereby applying the correction to $\Delta S_{\rm calc}^*$.

Table II contains a summary of electrochemical activation parameters for four Cr(III) reductions and also for ${\rm Eu}({\rm OH}_2)^{3+}_{n}$ reduction at mercury, lead, and upd lead surfaces. In addition to the values of ΔH_1^* determined from the temperature

The values of Γ_n for electrochemical reactions, Γ_n^e , as a function of temperature can be easily obtained from the quantities, Γ_n^h , derived from published relationships for homogeneous redox reactions by noting that only one redox center is activated in the former, versus two in the latter process. Thus for exchange reactions $\Gamma_n^e = (\Gamma_n^h)^{\frac{1}{2}}$. Similarly to Γ_n^h , Γ_n^e will decrease as the driving force (i.e. the overpotential) increases. For $\text{Cr}(\text{OH}_2)_6^{3+}$ reduction at -1000 mV, ΔS_1^* is calculated as -20 J K⁻¹ mol⁻¹, yielding $\Delta S_{\text{int}}^* \approx -13$ J K⁻¹ mol⁻¹ when this term is evaluated using the procedure given in ref. 39. A similar value of ΔS_{int}^* is likely for the other Cr(III) reactions considered here, even though exact calculations cannot be made. For $\text{Eu}(\text{OH}_2)_n^{3+}$ reduction, we estimate that $\Delta S_{\text{int}}^* \approx 0 \pm 5$ J deg⁻¹ mol⁻¹ in view of the smaller intrinsic barrier for this reaction together with smaller metalligand stretching frequencies.

These were determined using Eq. (7) from the listed values of ΔH_1^* and k_{corr}^* , along the composite preexponential factor $\delta r \kappa_{el} \Gamma_n \nu_n$. The last term was taken as 1.1 ± 10^5 cm sec⁻¹ for Cr(III) reductions; the constituent quantities $\Gamma_n = 2$, $\Gamma_n = 1.0 \times 10^{13} \text{ sec}^{-1}$ were obtained from the appropriate analytical expressions $\Gamma_n = 1.0 \times 10^{13} \text{ sec}^{-1}$ were obtained from the appropriate analytical expressions $\Gamma_n = 1.0 \times 10^{13} \text{ sec}^{-1}$ which the presumption that the "effective tunneling distance" $\Gamma_n = 1.0 \times 10^{13} \text{ sec}^{-1}$ is that expected if adiabaticity ($\Gamma_n = 1.0 \times 10^{13} \text{ sec}^{-1}$) is only achieved for the section sites at the plane of closest approach, $\Gamma_n = 1.0 \times 10^{13} \text{ sec}^{-1}$. The corresponding that the plane of closest approach, $\Gamma_n = 1.0 \times 10^{13} \text{ sec}^{-1}$. The corresponding that the plane of closest approach, $\Gamma_n = 1.0 \times 10^{13} \text{ sec}^{-1}$.

Two features are apparent upon examining the results in Table II. Most prominer in activation entropies for a given reaction became substantially smaller and even dependent upon substituting mercury by either lead or upd lead surfaces. However, where LS_1^* decreases are partially offset by decreases in the corresponding activation entralpies, especially for the aquo reactants $Cr(OH_2)_0^{3+}$ and $Eu(OH_2)_n^{3+}$. Also, in every case LS_1^* LS_1^* LS_1^* Although these values of LS_1^* specifically refer to the electrode potential -1000 mV, virtually the same values of LS_1^* (within experimental error, = 10-15 J K⁻¹ mol⁻¹) are obtained throughout the potential range where the table data were obtained. This latter result follows from the finding that the values were essentially independent of tempertaure, although small variations were detected for $Cr(NH_3)_0^{3+}$ reduction at mercury.

Though the temperature dependence of the double-layer corrections at the solid metals are known with less certainty than those at a given temperature due to the unanthown temperature dependence of the p.z.c., the resulting double-layer effect that the is liable to be small on the basis of such findings at mercury electrodes.

DISCUSSION

Dependence of Rate Parameters on Electrode Material

The kinetic data gathered in Tables I and II indicate that the energetics of most of these reactions depend upon the nature of the electrode material in a manner and to an extent which is clearly beyond that described by electrostatic double-layer effects. Particularly significant is the finding that the values of k for reactants containing largely aquo ligands decrease typically by factors up to 100-500 fold upon substituting liquid mercury by lead, upd thallium, and especially liquid gallium surfaces (Table I). Several factors may be envisaged as contributing to this finding on the basis of Eqs. (3) and (7). The simplest of these is the possibility that the effective tunneling distance $\delta r \kappa_{al}$ is decreased markedly upon altering the electrode material; i.e. the rate decreases are associated primarily with increasingly nonadiabatic reaction pathways. This is qualitatively consistent with the accompanying substantial decreases in ΔS_{i}^{*} especially since $\Delta S_{i}^{*} < \Delta S_{calc}^{*}$ (Table II). However, this explanation is not consistent with the concurrent decreases in ΔH_i^* that are observed, especially for the aquo reactants (Table II). This latter result suggests instead that the alterations in the electrode material produce substantial changes in the activation barrier to electron transfer.

These findings can be rationalized in terms of the varying influences of the metal surface upon the interfacial solvent structure. Evidence from a variety of sources suggests that the metal surfaces considered here vary widely in their tendency to orient inner-layer water molecules via metal-oxygen bonding. ^{26,27} Although some interpretations of measured physical properties along these lines evidently are fairly speculative, ⁴⁹ it is apparent that mercury has only a small tendency to crient water molecules in this manner, whereas gallium has a relatively strong

Tendency with lead and thallium being intermediate cases. ²⁷ Of the various such "hydrophilicity scales", that based on the enthalpy of formation, ΔH_{MO}° , of the appropriate bulk-phase metal oxide MO (or M₂O) seems to be relatively trustworthy. ⁴⁹ These data suggest that the order of hydrophilicity for the surfaces considered here is Hg < Pb ~ Tl < Ga. ²⁷ Interestingly, independent evidence supporting this assertion is obtained from infrared matrix isolation studies of adducts formed between metal atoms and water molecules. ²⁸ Thus adduct formation causes a systematic fecrease in the ν_2 bending mode of H₂O. Since the extent of this decrease, $\Delta \nu_2$, is fependent on the extent of σ metal-oxygen bonding, ²⁸ $\Delta \nu_2$ is anticipated to be related closely to the relative hydrophilicities of the corresponding metal surfaces. Indeed, an approximate correlation can be deduced between $\Delta \nu_2$ and ΔH_{MO}^{\bullet} for Group III and IV metals; this also yields the hydrophilicity order Pb \lesssim Tl < Ga.

Intriguing, the substantial substrate dependence of $k_{\rm corr}$ for the aquo meactants observed here largely falls in the same sequence, Hg < Pb < Ga (Table I). The behavior of the upd lead-silver surface is somewhat different to that of lead, the former generally yielding somewhat larger values of $k_{\rm corr}$ than the latter. Nevertheless, the $k_{\rm corr}$ values at upd Tl-Ag are generally somewhat smaller than those at upd Pb-Ag, in accordance with the above hydrophilicity order. On the other hand, the $k_{\rm corr}$ values for reduction of the two reactants not containing aquo ligands, ${\rm Cr(NH_3)}_6^{3+}$ and ${\rm Cr(en)}_3^{3+}$, show only a small dependence on the electrode material, although some variation in the activation parameters is observed (Tables I, II).

The aquo reactants therefore appear to display a particular sensitivity to the interfacial solvent environment. This recalls the finding of our earlier study 11 that the electrooxidation kinetics of several aquo reactants were decreased framatically at silver and especially platinum and gold relative to mercury. The

cuter-sphere reduction kinetics of Co(III) ammine complexes were found to be independent of the electrode material once electrostatic double-layer corrections were applied. While apparently similar, the present findings are more clear-cut since they refer to a series of structurally similar reactions at surfaces having well-defined, yet variable double-layer properties.

We have previously presented several lines of evidence indicating that transition-metal aquo reactants interact extensively with surrounding water molecules, i.e. are strongly hydrated. Thus although $\text{Cr}(\text{OH}_2)_6^{3+}$ and $\text{Cr}(\text{NH}_3)_6^{3+}$ have almost the same crystallographic radii, the former appears to undergo reduction at mercury some 1-2 Å further from the surface than the latter on the basis of detailed examinations of double-layer effects at this electrode 18,21,22 . This is consistent with the greater hydrated radii of $\text{Cr}(\text{OH}_2)_6^{3+}$ and other aquo complexes arising from strong ligand-solvent hydrogen bonding. The presence of such hydrogen-bonded secondary solvation also accounts for both the abnormally large reaction entropies 42 and large deuterium isotope effects upon the redox thermodynamics 50 and kinetics 20 of aquo redox couples. As noted previously, it is therefore anticipated that the solvation of aquo reactants would be unusually sensitive to differences between the surrounding solvent structure at the interface and the bulk solution. The occurence of a strong "surface hydrophilicity" effect for these reactants therefore seems reasonable.

Several consequences of this "surface solvent environmental" effect might be considered. Firstly, at hydrophilic surfaces the inner-layer solvent will tend to orient with the oxygen (or one oxygen lone pair) directed towards the surface. This ordering may propagate to the second solvent layer and even beyond. Such oriented solvent would find difficulty in hydrogen bonding to the incoming reactant, thereby significantly destabilizing the precursor state [i.e. yielding a smaller

 K_0 (or δr) Eq. (3)], and hence decreasing k_{corr} . Secondly, such a decrease in K_0 for sites suitably close to the surface will lead to the reaction occuring primarily via sites further from the metal surface, with smaller values of $\kappa_{\rm el}$. The interplay between these two factors should yield smaller effective tunneling distances (or K) and therefore smaller frequency factors, or, equivalently, smaller apparent activation entropies. A third, distinctly different, model invokes the interfacial potential drop, ϕ_{xy} , associated with water dipole orientation. Large negative values of ϕ_{xy} are deduced for hydrophilic surfaces, at least near the p.z.c. 27 The reactant may alter the interfacial solvent structure so to nullify this dipole orientation in its vicinity. The average value of $\boldsymbol{\varphi}_{_{\boldsymbol{U}}}$ across the surface should nonetheless be almost unaffected since the reactant surface coverage will be very small. Consequently, reactions at more hydrophilic surfaces will incur an additional overpotential, $\Delta \phi$, equal to ϕ_w , corresponding to a given value of k_{corr} . This explanation is at least roughly consistent with the data in Table I since these show that substantially (ca 300 mV) larger overpotentials are required to attain the same $k_{\mbox{corr}}$ values for aquo reactants at gallium compared with mercury; $\boldsymbol{\varphi}_{_{\!\!\boldsymbol{W}}}$ is estimated to be ca 500 mV for gallium at the p.z.c. 2/

These models do not, however, provide a ready explanation of the smaller activation enthalpies that generally accompany the decreases in k_{corr} seen at the more hydrophilic surfaces (Table II). A rationalization of this finding may nonetheless be made by postulating that the outer-sphere reactions occur at sites beyond the inner layer where local solvent "structure breaking" predominates. This invokes the "three-layer" model of interfacial solvent structure proposed by Drost-Hansen. ⁵¹ In this model, the strongly oriented water at a hydrophilic surface is separated from water possessing the normal bulk structure by an intermediate "structure-breaking" region where enhanced disorder occurs. This notation is akin to the secondary solvation region originally proposed in the Frank-Wen model of bulk ionic hydration. ⁵² Aque cations present within

such a "solvent-disordered" region would experience a smaller resistance to orienting surrounding water molecules than would be the case in bulk solution. This is due to the greater competition from solvent-solvent hydrogen bonding for the latter. Such enhanced solvent polarization would provide an enthalpic stabilization of the transition state, which would be offset by an entropic destabilization associated with solvent ordering. Although qualitative, this model accounts for the observed concurrent decreases in ΔH_1^* and ΔS_1^* as the hydrophilicity of the surface increases (Table II).

Although the values of α_{corr} for the various Cr(III) reductions vary noticeably with ligand composition, they do not significantly depend upon the electrode material (Table I). Consequently, the dependence of k_{corr} upon the metal surface

The same appears to be approximately true for the activation parameters, given the potential dependence of k_{corr} largely resides in the enthalpic component, This infers that the degree of hydrophilicity of these surfaces is not sensitive to the electrode potential, at least in the region where rate data could the obtained. Nevertheless, the more positive p.z.c. for mercury (-435 mV) compared thicse for the other metals considered here (-800 to -960 mV) may act to reinforce their differences in hydrophilicity. Thus the moderate negative charges $\frac{1}{2} \sim -8$ to $-12 \ \mu \text{C cm}^{-2}$) for the former surface at the electrode potentials where the rate data were obtained should further disfavor oxygen orientation towards the

Trisons with Theoretical Rate Parameters

The above interpretations suggest that the extent of the surface environmental influence upon the rate parameters is greatest for gallium and smallest for mercury. The conclusion can, in principle, be checked by comparing the individual rate Targeters directly with the numerical predictions from theoretical models of ===sition state experiences the same solvation environment as for the reactant === product states in the bulk solution. (This follows from the "weak overlap" Essemption noted above.) Under these circumstances the various contributions to === electron-transfer barrier arising from ion-solvent interactions will be accounted for providing that the electrochemical rate parameters are measured at a known thermo-====ic driving force. Consequently, one might expect that kinetic parameters messured at a mercury surface would be in closer accordance with the theoretical tradictions than those determined at a metal, such as gallium, that apparently tributes a large perturbation upon the interfacial solvent environment. (This presume of conventional electron-transfer theory are Interpretate, in particular the assumption that adiabatic pathways are followed.)

The comparison between "measured" and calculated activation entropies has ziready been considered. The advantage of this approach is that the theoretical zalculations of AS* do not require a detailed knowledge of the activation barrier. Erwever, it is clearly desirable to also compare the measured rate constants themserves with the theoretical predictions. This entails estimating the classical ectivation barrier, ΔG_{corr}^{\star} , as well as the preexponential factors in Eq. (3). For the present systems, ΔG_{corr}^{\star} includes large contributions from the inner-shell Earrier, Ω_{is}^* , associated with metal-ligand distortions, as well as the outer-shell tarrier, ΔG_{00}^{*} , arising from polarization of the surrounding solvent. Quantitative calculations of ΔG_{is}^* and hence k_{corr} are precluded for most of the present systems due to insufficient structural data. However, recent EXAFS measurements the chromium-aquo bond length difference, $\triangle a$, between $Cr(OH_2)_6^{3+}$ and $Cr(OH_2)_6^{2+}$ 53 sicng with values of the frequencies, v_3 and v_2 , of the Cr^{III}-OH, and Cr^{II}-OH, Eccas from vibrational spectroscopy 53,54 enable quantitative theoretical estimates of kcorr, k calc, to be obtained for this reaction. Details of these calculations are given elsewhere. 48,55 They employed the following parameters $= 2.0 \times 10^{-9}$ cm, 53 $v_3 = 540$ cm⁻¹, 54 and $v_2 = 380$ cm⁻¹, along with the usual Earmonic oscillator model for the inner shell, the dielectric continuum approximation for the outer shell (as in ref. 10) together with the above preexponential factors. These yield $k_{calc} \approx 5 \times 10^{-6}$ cm sec⁻¹ at the formal potential for $Cr(OH_2)_6^{3+/2+}$ (-655 mV^{14}) , and $k_{\text{calc}} \approx 5 \times 10^{-3} \text{ cm sec}^{-1} \text{ at } -1000 \text{ mV}$.

Considering the likely uncertainties in these calculations (at least tenfold $k_{\rm calc}$), the excellent agreement between $k_{\rm calc}$ at -1000 mV and $k_{\rm corr}$ at mercury $(3 \times 10^{-3} \ {\rm cm \ sec^{-1}})$, Table I) may be somewhat fortuitous. Nevertheless, this value $k_{\rm calc}$ is clearly much larger than the corresponding values of $k_{\rm corr}$ obtained at $k_{\rm calc}$ and especially gallium electrodes (Table I). This result therefore supports above assertion that the latter surfaces act to perturb structurally the $k_{\rm calc}$ above assertion state.

The activation parameter data for $\operatorname{Cr}(\operatorname{OH}_2)^{3+}_6$ reduction (Table II) are nicely sensistent with this picture. Thus the difference between ΔS_1^* and $\Delta S_{\operatorname{calc}}^*$, ΔK^{-1} mol⁻¹, only amounts to a ca tenfold discrepancy between the experimental calculated preexponential factors. This could be due in part to marginally calculated pathways, i.e. to "effective tunneling distances", $\delta r \kappa_{el}$, that are somewhat smaller than the value, $\delta \times 10^{-9}$ cm, assumed when evaluating $\Delta S_{\operatorname{calc}}^*$. Sowever, a similar value of $\delta r \kappa_{el}$ has been obtained for $\operatorname{Cr}(\operatorname{OH}_2)^{3+}_6$ reduction at mercury from a comparison of experimental inner—and outer—sphere reactivities. Elternatively, this decrease could be due to additional solvent ordering in the transition state caused by the presence of the electrode surface. The much larger differences between ΔS_1^* and $\Delta S_{\operatorname{calc}}^*$ seem at the more hydrophilic surfaces (Table II) seem more consistent with this latter interpretation.

A similar trend is also seen for $\mathrm{Eu}(\mathrm{OH}_2)_n^{3+}$ reduction, the difference between -5 and $\Delta S_{\mathrm{calc}}^*$ at mercury also being much smaller than at lead and upd lead-silver, elthough the former difference (52 J K⁻¹ mol⁻¹) corresponds to a 500-fold discrepancy -1 the preexponential factor. This may be associated in part with the occurrence of -1 to -1 to -1 the preexponential factor. This may be associated in part with the occurrence of -1 to -1 the preexponential factor -1 the preexponential factor -1 to -1 the preexponential factor -1 the pr

If nonadiabaticity provides an important contribution to the measured substrate effects, one factor influencing the values of κ_{el} would be the electron density distribution at the metal surface. This distribution is expected to be sensitive to the chemical nature of the metal; lead and especially gallium should have disher electron densities than mercury, protruding further from the metal surface. This factor should therefore yield enhanced electronic coupling with the reactant acceptor orbitals, and hence larger values of κ_{el} and κ_{corr} lead and gallium relative to mercury, in complete contrast to the experimental

CONCLUSIONS

The present results, together with those from our earlier study. 11 attest to The important influence that the metal surface can exert upon the energetics of even outer-sphere electrochemical reactions involving strongly hydrated reactants. As noted previously, 11 it appears useful to divide outer-sphere reactions into "solvent structure-demanding" and "structure-undemanding" categories, depending on the sensitivity of the reaction energetics to the interfacial reaction environment. It is possible that the cationic aquo complexes considered here are unusually, even uniquely, "solvent structure-demanding" reactants in that especially large dependences of k_{corr} on the nature of the electrode material are obtained. Nevertheless, other systems may also display a strong sensitivity of the enthalpic and entropic components of the activation barrier to the interfacial environment, but so that these components largely cancel, yielding relatively structure-independent electrochemical reactivities. Such enthalpy-entropy compensation is a well-known phenomenon. especially for aqueous systems.⁵⁸ The reduction of $Cr(NH_3)_6^{3+}$ examined here appears to be such an example. Substantially different behavior is also anticipated in nonaqueous media, especially in solvents for which hydrogen bonding or other strong intermolecular interactions are absent. 55

It is also interesting to relate the present electrochemical results to those involving outer-sphere reactions in homogeneous aqueous solution between structurally similar transition-metal reactants to those considered here. 46,48,55 Broadly speaking the energetics of such reactions involving complexes containing aquo, ammine, polypyridine or related ligands show similar deviations from the theoretical expectations as are observed for the present electrochemical reactions at hydrophilic metal surfaces. Thus generally $k_{\rm calc} < k_{\rm corr}$, and $\Delta S^* << \Delta S^*_{\rm calc}$. These deviations, which are somewhat dependent upon the ligand structure of the

coreactant, can be attributed in part to marginally nonadiabatic pathways $(\kappa_{el} \ge 0.01)^{53}$ and the entropically unfavorable perturbation of the local solvent structure induced by the nearby coreactant. Therefore at least some vagaries of the structure-sensitive energetics of electrochemical and homogeneous outer-sphere reactions may well have common origins.

ACKNOWLEDGMENTS

This work is supported in part by the Air Force Office of Scientific Research and the Office of Naval Research. M.J.W. acknowledges a fellowship from the Alfred P. Sloan Foundation.

REFERENCES

- 1. J. T. Hupp, M. J. Weaver, J. Electroanal. Chem. 152 (1983), 1.
- 2. N. Sutin, B. S. Brunschwig, ACS Symp. Ser. 198 (1982), 105.
- 3. S. W. Barr, N. J. Weaver, Inorg. Chem., in press; K. L. Guyer, M. J. Weaver, Inorg. Chem., in press.
- -. S. W. Barr, K. L. Guyer, T. T-T. Li, H. Y. Liu, M. J. Weaver, J. Electrochem. Soc., <u>131</u> (1984), 1626.
- 5. S. Trassati, in "Advances in Electrochemistry and Electrochemical Engineering", H. Gerischer, C. W. Tobias (eds), Wiley, New York, Vol. X, 1977, p. 279.
- 6. P. Delahay, "Double Layer and Electrode Kinetics", Interscience, New York, 1965, Chapter 9.
- 7. M. J. Weaver, J. Electroanal. Chem. 93 (1978), 231.
- 8. W. R. Fawcett, "Proc. 3rd Symp. on Electrode Processes", S. Bruckenstein, B. Miller, J. D. E. McIntyre, E. Yeager (eds), Electrochemical Society, Pennington, N.J., 1980, p. 213; M. W. Weaver, *ibid*, p. 233.
- 9. R. A. Marcus, Can. J. Chem. 37 (1959), 155.
- _O. M. J. Weaver, J. Phys. Chem. 84 (1980), 568.
- 1. S. W. Barr, K. L. Guyer, M. J. Weaver, J. Electroanal. Chem. 111 (1980), 41.
- 2. A. Capon, R. Parsons, J. Electroanal. Chem. 46 (1973), 215.
- 13. For example, A. N. Frumkin, N. V. Fedorovich, S. I. Kulakoskaya, Sov. Electrochem. 10 (1973), 313.
- 14. M. J. Weaver, F. C. Anson, J. Electroanal. Chem. 65 (1975), 711.
- 15. M. J. Weaver, F. C. Anson, J. Electroanal. Chem. 65 (1975), 737, 759.
- L6. M. J. Weaver, F. C. Anson, Inorg. Chem. 15 (1976), 1871.
- 17. M. J. Weaver, F. C. Anson, J. Phys. Chem. 80 (1976), 1861.
- 18. M. J. Weaver, T. L. Satterberg, J. Phys. Chem. 81 (1977), 1772.
- 19. M. J. Weaver, J. Phys. Chem. 83 (1979), 1748.
- 20. M. J. Weaver, P. D. Tyma, S. M. Nettles, J. Electroanal. Chem. 114 (1980), 53.
- 11. M. J. Weaver, H. Y. Liu, Y. Kim, Can. J. Chem. 59 (1981), 1944.
- _2. J. T. Hupp, M. J. Weaver, J. Phys. Chem., 88 (1984), 1463.
- _3. H. Y. Liu, Ph.D. thesis, Michigan State University, 1982.

- 1-. J. T. Hupp, Ph.D. thesis, Michigan State University, 1983.
- 15. J. T. Hupp, D. Larkin, H. Y. Liu, M. J. Weaver, J. Electroanal. Chem. <u>131</u> (1982), 299.
- 25. B. B. Damaskin, A. N. Frumkin, Electrochim. Acta 19 (1974), 173.
- 17. S. Trassati, "Modern Aspects of Electrochemistry", Vol. 13. B. E. Conway, J. O'M. Bockris (eds), Plenum, New York, 1979, p. 81; S. Trassati, Electrochim. Acta. 28 (1983), 1083.
- 28. R. H. Hauge, J. W. Kauffman, L. Fredin, J. L. Margrave, ACS Symp. Ser. 179 (1982), 363; R. H. Hauge, J. W. Kauffman, J. L. Margrave, J. Am. Chem. Soc. 102 (1980), 6005.
- 29. K. L. Guyer, Ph.D. thesis, Michigan State University, 1981.
- 3C. A. Bewick, J. Robinson, J. Electroanal. Chem. 60 (1975), 163.
- 31. J. P. Carr, N. A. Hampson, R. Taylor, J. Electroanal. Chem. 32 (1971), 345.
- 31. K. B. Oldham, E. P. Parry, Anal. Chem. 40 (1968), 65.
- 33. A. J. Bard, L. R. Faulkner, "Electrochemical Reactions", Wiley, New York, 1980, p. 223.
- 5-. D. Larkin, K. L. Guyer, J. T. Hupp, M. J. Weaver, J. Electroanal. Chem. <u>138</u> (1982) 401.
- 35. I. A. Bagostskaya, A. M. Morozov, N. B. Grigoryev, Electrochim. Acta. 13 (1968), 873.
- 35. J. T. Hupp, H. Y. Liu, M. J. Weaver, in preparation.
- 37. K. L. Guyer, S. W. Barr, R. J. Cave, M. J. Weaver in "Proc. 3rd Symp. on Electrode Processes", S. Bruckenstein, J. D. E. McIntyre, B. Miller, E. Yeager (eds), Electrochemical Society, Pennington, 1980, p. 390.
- 35. A. N. Frumkin, N. B. Grigorev, I. A. Bagotskaya, Dokl. Akad. Nauk SSSR, 157 (1964), 957; V. F. Ivanov, Z. N. Ushakova, Sov. Electrochim. 9 (1973), 753.
- 35. J. T. Hupp, M. J. Weaver, J. Phys. Chem., 88 (1984), 1860.
- -1. M. J. Weaver, J. Phys. Chem. 80 (1976), 2645.
- -I. J. T. Hupp, M. J. Weaver, J. Electroanal. Chem. 145 (1983), 43.
- -I. E. L. Yee, R. J. Cave, K. L. Guyer, P. D. Tyma, M. J. Weaver, J. Am. Chem. Soc. 101 (1979), 1131.
- -1. S. Sahami, M. J. Weaver, J. Electroanal. Chem. 122 (1981), 155, 171.
- --. J. T. Hupp, M. J. Weaver, Inorg. Chem., in press.
- +1. 5. S. Brunschwig, J. Logan, M. D. Newton, N. Sutin, J. Am. Chem. Soc. 102 (1980), 5798.

- 46. N. Sutin, Prog. Inorg. Chem. 30 (1983), 441.
- 47. E. L. Yee, J. T. Hupp, M. J. Weaver, Inorg. Chem. 22 (1983), 3465.
- 48. J. T. Hupp, M. J. Weaver, submitted for publication.
- 49. G. Valette, J. Electroanal. Chem. 139 (1982), 285.
- 50. M. J. Weaver, S. M. Nettles, Inorg. Chem. 19 (1980), 1641.
- 51. W. Drost-Hansen, Ind. Eng. Chem. 61 (11) (1969), 10.
- 52. H. S. Frank, W. Y. Wen, Disc. Far. Soc. 24 (1957), 133.
- 53. B. S. Brunschwig, C. Creutz, D. H. McCartney, T-K. Sham, N. Sutin, Disc. Far. Soc. 74 (1982), 113.
- 54. S. P. Best, J. K. Beattie, R. S. Armstrong, J. Chem. Soc. Dalton, in press; T. E. Jenkins, J. Lewis, Spectrochim. Acta 37A (1981), 47.
- 55. J. T. Hupp, H. Y. Liu, J. K. Farmer, T. Gennett, M. J. Weaver, J. Electroanal. Chem., in press; presented at the Sixth Australian Conference on Electrochemistry, Geelong, Victoria, February 1984.
- J. P. Badiali, M. L. Rosinberg, J. Goodisman, J. Electroanal. Chem. <u>130</u> (1981), 31; *ibid*, <u>143</u> (1983), 73.
- 57. N. D. Lang, Solid State Phys. 28 (1973), 225.
- 58. R. Lumry, S. Rajander, Biopolymers 9 (1976), 1125.
- 59. M. J. Weaver, E. L. Yee, Inorg. Chem. 19 (1980), 1936.

TABLE I. Rate Constants for the Electroreduction of Cr(III) and related

Trivalent Complexes at -1000 mV vs s.c.e. at Various Metal Surfaces
at 25°C.

Reactant	Surface	k a app cm sec ⁻¹	app b	k c corr cm sec 1	α d corr
Cr(OH ₂) ₆ 3+	Нg ~	5 x 10 ⁻²	0.61	3 x 10 ⁻³	0.50
	Ga	8×10^{-6}	0.58	3×10^{-6}	0.50
4	Pb	3×10^{-5}	0.61	1.5×10^{-5}	0.55
	upd Pb-Ag	3×10^{-3}	0.55	1.5×10^{-3}	0.52
	upd Tl-Ag	3×10^{-4}	0.50	3×10^{-4}	0.50
Cr(OH ₂) ₅ F ²⁺	Hg	2.5×10^{-4}	0.58	2.5×10^{-5}	0.54
	Pb	2×10^{-6}	0.55	1×10^{-6}	0.52
	upd Pb-Ag	$^{-2} \times 10^{-5}$	0.55±0.05	$\sim 1 \times 10^{-5}$	~0.55
	upd Tl-Ag	1.5×10^{-6}	0.65	1.0×10^{-6}	0.6
Cr(OH ₂) ₅ OSO ₃ +	Нg	3.5×10^{-3}	0.54	2 x 10 ⁻³	0.52
	Ga	$^{-7} \times 10^{-6}$	~0.55±0.05	$\sim 5 \times 10^{-6}$	≈0.55
	Pb	3×10^{-5}	0.5±0.05	2.5×10^{-5}	≈0.5
	upd Pb-Ag	5 x 10 ⁻⁵	0.5±0.05	3×10^{-5}	≈0.5
Cr(NH ₃) ₆ ³⁺	Hg	2 x 10 ⁻²	0.84	4 x 10 ⁻⁴	0.75
	Ga	~2 x 10 ⁻⁴	0.8±0.05	1.5×10^{-4}	≈0. 7
	Pb	6×10^{-6}	0.76	1.5×10^{-6}	0.7
	upd Pb-Ag	2.5×10^{-4}	0.78	7 x 10 ⁻⁵	0.70
	upd Ti-Ag	8×10^{-5}	0.70	7×10^{-5}	0.65

Cr(NH ₃)50H ₂ 3+	Нg	6×10^{-2}	0.75	2 × 10 ⁻³
	Ga	3×10^{-5}	0.70	2 × 10 ⁻⁵
	Pb	2 x 10 ⁻⁵	0.73	5 × 10 ⁻⁶
	upd Pb-Ag	1 x 10 ⁻³	0.8	4 × 10 ⁻⁴
	upd Tl-Ag	9×10^{-5}	0.7	e × 10 ⁻⁵
Cr(en)3+ e	Hg	7 x 10 ⁻²	0.90	3 × 10 ⁻⁴
	upd Pb-Ag	1×10^{-3}	0.90	5 × 10 ⁻⁴
Eu(OH ₂) ^{3+ f}	Hg	~4	0.62	ر, 1
2 11	Pb	6×10^{-3}	0.41	$_{2}$ $_{10}^{-3}$
	upd Pb-Ag	2×10^{-2}	0.41	1 × 10 ⁻²
v(он ₂) ₆ 3+	Нg	~20	0.4-0.5	~1,.5
2′6	**6		0.4-0.5	
	Pb	0.2	0.45	5 × 10 ⁻²

Apparent (i.e. observed) rate constant for one-electron electron eduction at electrode potential E = -1000 mV at metal surface listed; electroly NaClO₄ + 3 mM HClO₄, except for Cr(NH₃) $_{3}^{3+}$ reduction which was mossured La(ClO₄) $_{3}$ + $\overline{3}$ mM HClO₄. Reproducibility of k_{app} generally $\pm 10\%$ at merotate other surfaces. Values of k_{app} below ca. 1×10^{-4} cm sec⁻¹ extrapoly values measured at more negative potentials from (Tafel) plot of k_{app}

b Apparent transfer coefficient, determined from $\alpha_{\rm app} = -({\rm RT/F}) ({\rm d} i)^{ik} {\rm app}/i$ and measurement conditions as indicated for corresponding value: of kallisted are average quantities determined at electrode potential corresponding values from ca. 10^{-4} to 10^{-2} cm sec⁻¹. Reproducibility of $\alpha_{\rm app}$ mercury and ± 0.02 at other surfaces except where indicated.

Rate constant at -1000 mV corrected for electrostatic work term; , determined value of k using Eq. (2); required values of the diffire-lay, determined as noted in text and in footnotes below.

dTransfer coefficient corrected for electrostatic work terms, de, ermine, value of α_{app} using Eq. (6). Required values of $(d\phi_r/dE)$ determined for Cd1/Cdiff, where Cd1 is the measured double-layer capacitance of Cdiff layer capacitance. For 0.5 M NaClO4, Cdiff from 0.6 Cdiff CGC where Gouy-Chapman estimate; for 40 mM La(ClO4)3, from Cdiff CGC See 1

en = ethylenediamine

fn denotes unknown number (probably 8-9) of aquo ligands.

THE II. Electrochemical Activation Parameters for the Electroreduction of Cr(III) and Eu(III) Reactants at -1000 mV vs s.c.e. at Mercury, Lead, and upd Lead Surfaces.

F.esctant	Surface	k corr cm sec ⁻¹		∆S* c J K ⁻¹ mol ⁻¹	J K ⁻¹ m
C= (OH ₂) ₆ ³⁺	Hg	3 x 10 ⁻³	63	68	 87 ^{e}
	Pb	1.5×10^{-5}	48	-27	87 ^e
	upd Pb-Ag	1.5×10^{-3}	30	-49	87 [€]
C= (NH ₃) ³⁺	Нg	4 x 10 ⁻⁴	57	30	(38) ^f
•	Pb	1.5×10^{-6}	44	-60	(38) ^f
	upd Pb-Ag	7×10^{-5}	45	-25	(38) ^f
C= NH ₃) ₅ OH ₂ ³⁺	Hg	2 x 10 ⁻³	56	40	
	Pb	5×10^{-6}	31	-94	
	upd Pb-Ag	4 x 10 ⁻⁴	58	34	• • •
C r (en) 3+	Hg	3×10^{-4}	62 · ·	44	
	upd Pb-Ag	5 x 10 ⁻⁴	55	25	• • •
≣ਦ (0H ₂) ³⁺	Hg	0,2	48	48	100 ^g
- **	РЪ	2.5×10^{-3}	31	-45	70 ⁸
	upd Pb-Ag	1 x 10 ⁻²	24	-57	70 ⁸

Footnotes to Table II

Work-corrected rate constant for reduction of complex at surface indicated at -1000 mV; taken from Table I.

b"Ideal" enthalpy of activation at -1000 mV; determined from $\Delta H_2^* = -R[d\ln k_{\text{COTT}}/d(1/T)]$ using a nonisothermal cell arrangement with the saturated calomel reference electrode held at room temperature. Reproducibilities of ΔH_2^* are: mercury, \pm 1 kJ mol⁻¹; solid electrodes, \pm 5 kJ mol⁻¹, except for $Cr(NH_3)_6^{2+}$ reduction at upd lead (\pm 15 kJ mol⁻¹), and $Cr(NH_3)_5OH_2^{2+}$ reduction at upd lead (\pm 10 kJ mol⁻¹).

C"Ideal" entropy of activation at -1000 mV, obtained from Eq. (7); i.e. from $\Delta S^* = R(\ln k - \ln A + \Delta H^*/RT)$, where $A(=\delta r \kappa \Gamma \nu)$ taken as $1.0^{\circ} \times 10^{\circ}$ cm $^{\circ}$ sec $^{\circ}$ for Cr(III) reactants and $^{\circ}$ $^{\circ}$ $^{\circ}$ $^{\circ}$ $^{\circ}$ 104 cm sec $^{\circ}$ for Eu(III). (See text for details.)

^dCalculated entropy of activation, obtained from Eq. (8) using α as listed in Table I, and values of ΔS_{int}^* and ΔS_{rc}° as indicated in footnotes below.

 $^{\rm e}$ Using $\Delta S_{\rm int}^{\star} = -13$ J K $^{-1}$ mol $^{-1}$ (see text), and $\Delta S_{\rm rc}^{\circ} = 205$ J K $^{-1}$ mol $^{-1}$. 42

fusing $\Delta S_{int_3+/2+}^* = -13 \text{ J K}^{-1} \text{ mol}^{-1}$ (see text), and ΔS_{rc}^* assumed to equal that for $Ru(NH_2)_6^{3+/2+}$ couple (75.5 J K⁻¹ mol⁻¹ ⁴²).

 $g_{\text{Using }\Delta S_{\text{int}}^{*}} = 0$ (see text), and $\Delta S_{\text{rc}}^{*} = 200 \text{ J K}^{-1} \text{ mol}^{-1}.42$

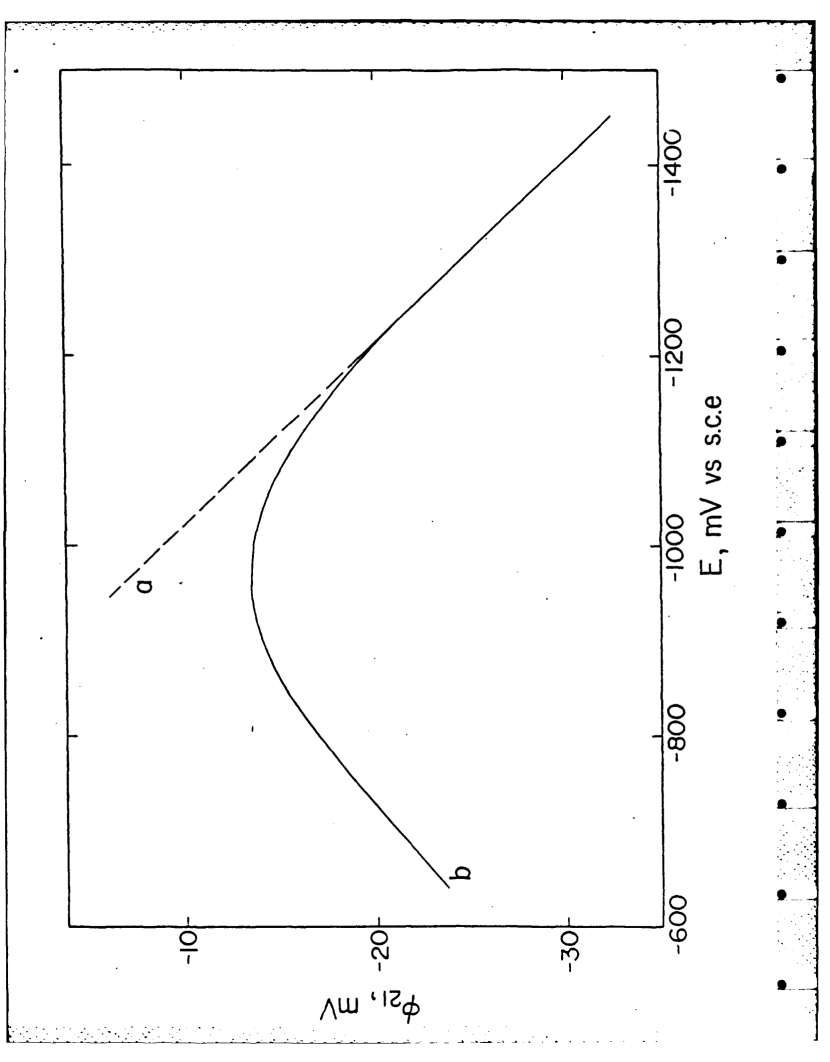


Figure Caption

Plot of diffuse-layer potential ϕ_2 against the electrode potential E for (a) 0.5 M NaF and (b) 0.5 M NaClO4 aqueous electrolytes in contact with a upd lead-silver surface. Values of ϕ_2 obtained using Gouy-Chapman theory and double-layer compositional data extracted from differential capacitance-potential measurements as outlined in the text.

Intripriate, in particular the assumption that adiabatic pathways are followed.)

END

FILMED

10-84

DTIC